UC Berkeley

UC Berkeley Previously Published Works

Title

Radiochemical Separation Techniques in Classroom Settings

Permalink

https://escholarship.org/uc/item/6516w4pj

Journal

Journal of Chemical Education, 100(4)

ISSN

0021-9584

Authors

Wang, Yufei

Abergel, Rebecca J

Publication Date

2023-04-11

DOI

10.1021/acs.jchemed.2c01193

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at https://creativecommons.org/licenses/by/4.0/

Peer reviewed





Laboratory Experiment



pubs.acs.org/jchemeduc

Radiochemical Separation Techniques in Classroom Settings

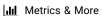
Yufei Wang* and Rebecca J. Abergel

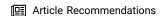


Cite This: J. Chem. Educ. 2023, 100, 1603-1612



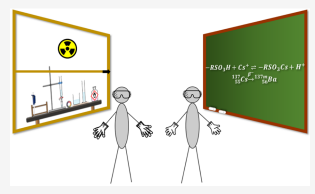
ACCESS I







ABSTRACT: In view of the mounting requirements for courses in nuclear chemistry and radiochemistry, hands-on experience in the separations of radioisotopes is imperative. Three separation techniques commonly used were introduced, coprecipitation, liquid-liquid extraction, and ion-exchange chromatography. Their principles and the associated fundamental thermodynamic concepts are detailed. Their respective experimental applications to the separations of radioisotopes of one rare earth element (europium) and two transition metals (cobalt and nickel) are discussed. In-classroom experimental settings are proposed, and experiment outcomes are presented, which can be beneficial for future implementation within nuclear chemistry and radiochemistry programs in higher education.



KEYWORDS: Upper-Division Undergraduate, Interdisciplinary/Multidisciplinary, Hands-On Learning/Manipulatives, Nuclear Chemistry/Radiochemistry, Separation Science

1. INTRODUCTION

The past few years have witnessed increased needs for a workforce in nuclear chemistry and radiochemistry (N&RC) expertise due to sustained efforts in radiopharmaceutical development, medical imaging and radiation oncology technology renewal, nuclear power production, waste management operations, and several other industrial sectors. Unfortunately, there is an unequivocal training and recruitment crisis that has led to a widening gap between a retiring workforce and early career radiochemists in many countries including the United States. This is partly attributable to the fact that the United States currently offers only a handful of undergraduate and graduate programs in N&RC. The number of people with N&RC expertise has stagnated at a low level, if not steadily decreased, over the past five decades in the United States, although N&RC-pertaining programs (e.g., chemistry, chemical engineering, and nuclear engineering) are prevalent. In contrast, China is experiencing a renaissance in these fields because of a very large demand for nuclear energy, and many academic programs have been established to meet this demand including both undergraduate and graduate-level degrees in radiochemistry and nuclear chemical engineering.³ To fill this gap in the United States, degree-granting programs, or courses at the minimum, in N&RC ought to be widely accessible. The utilization of virtual reality technology has enabled an unprecedented education mode and such application has been attempted in teaching introductory N&RC courses, too.^{4,5} This is particularly meaningful during the COVID-19 pandemic. However, the N&RC profession requires the mastery of hands-on experience, so the simulated experiments may assist but can never replace hands-on experiments. "Real"

N&RC experiments often use either sealed sources to detect radiation^{6,7} or surrogates in radiometric analysis^{8,9} to avoid direct handling of dispersive radioactive materials. Working with dispersive radioactive materials is a critical component in the N&RC profession, especially in radiochemical separation.

Benefiting from a well-established research ecosystem in nuclear and chemical sciences, the University of California at Berkeley (UC Berkeley) has long sought to familiarize students with N&RC principles and to provide hands-on training for the safe handling of radioactive materials in the laboratory. To that end, the Radiochemical Methods in Nuclear Technology and Forensics course (CHEM/NE C146) is offered to upperdivision undergraduate students and cross-listed between the Departments of Chemistry and Nuclear Engineering. The topics covered in this course were organized largely in accordance with the curriculum created over half a century ago, 10,11 starting with radiation detection and progressing to wet chemistry involving the manipulation of radioisotopes. The main distinction is that the CHEM/NE C146 course decreased the introduction of certain types of detectors (e.g., β -ray counters) and increased the number of wet chemistry experiments (e.g., uranium electrodeposition for nuclear forensics application). Among the numerous radiochemical

Received: December 8, 2022 Revised: February 25, 2023 Published: March 16, 2023





procedures investigated to date, including isotope dilution analysis, 12,13 radiochemical assay, 14 and radiochemical separation, the latter stands out due to its diverse applications in many nuclear sectors such as the production and purification of medical isotopes, the fabrication of nuclear fuel components, and the recovery of fission products and actinides for environmental remediation and strategic purposes. For training purposes, it is crucial to design radiochemical separation experiments that are easy and safe to perform, effective in providing experience in handling dispersive radioactive materials, and accessible to most universities and colleges. Here we aim to disseminate the UC Berkeley course's approach and experimental setups that have been refined over the years so that they can be implemented at other institutions. This work therefore starts with enunciating a few fundamental thermodynamic concepts in separation processes before introducing three separation techniques commonly used in N&RC, coprecipitation (Co-ppt), liquid-liquid extraction (LLE, also known as solvent extraction), and ion-exchange chromatography (IEC). The exemplary radioisotopes to be separated are europium-152 (152Eu3+), cobalt-60 (60Co2+), and nickel-63 (63Ni²⁺) (charges of metal ions will be omitted for simplicity from here on). Herein, experimental designs for each technique are proposed, experiment results presented, and possible adjustments discussed.

2. FUNDAMENTAL THERMODYNAMICS CONCEPTS IN SEPARATION PROCESSES

A separating agent (either mass or energy) is added into a mixture and subsequent dynamic separation of target species usually relies on either mechanical forces (e.g., filtration, centrifugation) or differences in mass transfer. When differences in mass transfer occur in different phases, the separation process relies on different thermodynamic equilibria (e.g., liquid-liquid extraction, chromatography, distillation). In contrast, when differences in mass transfer occur in the same phase, the separation process relies on different mass transfer rates (e.g., gas diffusion, reverse osmosis, electrodialysis). The three radiochemical separation techniques discussed here (Coppt, LLE, and IEC) are all based on differences in thermodynamic equilibria of substances and are commonly used in the classroom, research laboratory, and industrial settings. Familiarization with these three separation techniques will benefit education and training in N&RC. Separation processes involve aspects of thermodynamics, kinetics, and transport processes (momentum, energy (heat), and mass), with concepts in thermodynamics being most essential. Kinetics and transport processes are beyond the scope of this work, and a variety of textbooks can be referred to for further interest.

2.1. Partition Constant (K°), Partition Ratio (K), and Distribution Ratio (D)

One common confusion that many students have is among partition constant (K°) , partition ratio (K), and distribution ratio (D). Although they are often used interchangeably, the IUPAC Orange Book has made distinguished use of each. Both K° and K refer to the distribution of a particular chemical species with a definite form (e.g., a metal, denoted by M) across two distinct phases (eq 1), where the two phases for each of the three separation techniques are summarized in Table 1. However, these two parameters are inherently different. Considering a simple physical distribution of M

Table 1. Phases Assigned in Different Radiochemical Separation Techniques

	Co-Precipitation	Liquid—Liquid Extraction	Ion-Exchange Chromatography
Phase 1	Filtrate	Aqueous	Mobile
Phase 2	Precipitate or Filtrant	Organic	Stationary

across two phases without chemical reactions, K° can be derived from the phase equilibrium criterion and correlated to K via the equilibrium constant of the distribution reaction (eq 2). K° is independent of phase composition and dependent on temperature exclusively. In contrast, K, defined as $K = C_{\text{pha2}}/$ C_{pha1} (Nernst's distribution law), depends on both composition and temperature and is measured under specific conditions. Another difference is that K° is unitless, whereas K can be unitless or with unit, depending on whether the concentrations across two phases are expressed in the same unit (e.g., in LLE) or different units (e.g., in IEC where mol of M/g stationary phase and mol of M/mL mobile phase are commonly used). It is necessary to reiterate that even if they are approximately equal, as in many radiochemistry studies where solutions are sufficiently dilute, they are not identical. In the interest of metal separations, M often exists as multiple species in a phase (e.g., different complexed forms, different oxidation states, etc.), rendering K° and K inapplicable and necessitating the use of another thermodynamic variable, *D*:

$$M_{pha1} \rightleftharpoons M_{pha2} \qquad (1)$$

$$\underline{\mu_{pha1}^{\circ} + RT \ln a_{pha1}} = \underline{\mu_{pha2}^{\circ} + RT \ln a_{pha2}} \Rightarrow K^{\circ} = \frac{a_{pha2}}{a_{pha1}}$$

$$= \exp\left(-\frac{\mu_{pha2}^{\circ} - \mu_{pha1}^{\circ}}{RT}\right) = \operatorname{const}_{T} \stackrel{a=\gamma C}{\Longrightarrow} K^{\circ}$$

$$= \underbrace{\left(\frac{C_{pha2}}{C_{pha1}}\right)}_{K} \underbrace{\gamma_{pha1}}_{\gamma_{pha1}} \Rightarrow K = \operatorname{const}_{T,\gamma_{pha2}/\gamma_{pha1}} \qquad (2)$$

where μ_{pha} , a_{pha} , C_{pha} , and γ_{pha} are, respectively, the chemical potential, activity, concentration, and activity coefficient in a phase; R, universal gas constant; T, absolute temperature.

In contrast to K° and K, the distribution ratio D refers to the total analytical concentration of a solute in two phases, regardless of the chemical form (whether dissociated, associated, hydrolyzed, or complexed) (eq 3) and is dependent on both temperature and phase composition:

$$D = \frac{C_{tot,pha2}}{C_{tot,pha1}} = \frac{\sum_{i} [M_{i}]_{pha2}}{\sum_{j} [M_{j}]_{pha1}}$$
(3)

D, like K, can also be unitless or with unit, depending on the unit expression of concentration. When the solute only exists in one form, D and K are equal. Otherwise, D and K are correlated by the equilibrium constant of a reaction (dissociation, association, hydrolysis, or complexation) that produces the species. For instance, when a metal ion is complexed by a ligand (L) and forms a complex (ML_n , where $n = 1, 2, 3, \ldots$), D can be expressed as a function of K and $\log \beta$, where β is the stability constant of the complex.

Table 2. Separation Performance Evaluation Parameters Used in This Work

	Co-ppt	LLE	IEC	
Reaction Equation	$M_{(liq)} + c_{(p)} \rightleftharpoons M_{(p)} + c_{(liq)}$ where c represents carrier.	$M_{(aq)} \rightleftharpoons M_{(org)}$	CEC: $n\overline{RH} + M^{n+} \rightleftharpoons \overline{R_nM} + nH^+$ AEC: $m\overline{ROH} + ML_x^{m-} \rightleftharpoons \overline{R_mML_x} + mOH^-$ where $\overline{}$ represents resin.	
Selectivity coefficient, $K_{A/B}$	$K_{M/c} = \frac{K_M}{K_c} = \frac{n_{M,p}/n_{M,ml}}{n_{c,p}/n_{c,ml}}$ a	Often misused for SF, not recommended by IUPAC	$K_{M/(nH)} = \frac{K_M}{K_H^n}$	
$SF, SF_{M_1/M_2} = \frac{\kappa_{M_1}}{\kappa_{M_2}} b$	$SF_{M_1/M_2} = \frac{K_{M_1/c}}{K_{M_2/c}} = \frac{n_{M_1,p}/n_{M_1,ml}}{n_{M_2,p}/n_{M_2,ml}}$	$SF_{M_1/M_2} = \frac{C_{M_1,o}/C_{M_1,a}}{C_{M_2,o}/C_{M_2,a}}$	$SF_{M_1/M_2} = \frac{\kappa_{M_1/(n_1H)} \ \kappa_H^{n_1}}{\kappa_{M_2/(n_2H)} \ \kappa_H^{n_2}} \circ$	
Recovery, $R = \frac{Q_{pha2}}{\frac{Q_{pha1} + Q_{pha2}}{Q_0}}$	$R = \frac{n_p}{CV + n_p}$	$R = \frac{C_o V_o}{C_o V_o + C_a V_a} = \frac{D}{D + 1/\alpha}$	$R = \frac{C_e V_e}{C_S V_S + C_e V_e} = \frac{D}{D + 1/\alpha}$	
EF, $EF_{M_1/M_2} = \frac{R_{M_1}}{R_{M_2}} = \frac{Q_{M_1}/Q_{M_2}}{Q_{M_1,0}/Q_{M_2,0}} d$	$EF_{M_1/M_2} = \frac{n_{M_1,p}/n_{M_2,p}}{C_{M_1,0}/C_{M_2,0}}$	$EF_{M_1/M_2} = \frac{D_{M_1}}{D_{M_2}} \frac{D_{M_2} + 1/\alpha}{D_{M_1} + 1/\alpha}$	$EF_{M_1/M_2} = \frac{D_{M_1}}{D_{M_2}} \frac{D_{M_2} + 1/\alpha}{D_{M_1} + 1/\alpha}$	

^aFor linear/homogeneous distribution, the Berthelot-Nernst distribution law applies. ^bAssume single species of M_1 and M_2 . Replace K with D if M_1 and M_2 have multiple chemical forms. ^cSF in IEC can also be calculated by relative retention. $RR_{i,ref} = \frac{k_i}{k_{ref}} = \frac{K_i}{K_{ref}} = \frac{(x_i'/x_m)_i}{(x_i'/x_m)_{ref}}$, where x is t or

$$V \Rightarrow \mathit{SF}_{\mathit{M}_{1}/\mathit{M}_{2}} = \frac{\mathit{RR}_{1,\mathit{ref}}}{\mathit{RR}_{2,\mathit{ref}}}.~^{\mathit{d}}\mathit{EF}_{\mathit{M}_{1}/\mathit{M}_{2}} = \frac{1}{\mathit{DF}_{\mathit{M}_{1}/\mathit{M}_{2}}}.$$

In radiochemistry, the radioisotope concentration is often too low for chemical analysis to accurately quantify, and radiation detectors are most often resorted to. Detection of radiations usually gives total counts or count rates in counts per unit time that can be converted to radioactivity, from which the isotope concentration can be calculated. Since decay properties are not influenced by the phase in which the radionuclide exists and phase volume ratios are specified prior to the experiment, measuring the ratio of radioactivity between two phases is enough to ascertain D (eq 4):

$$D = \frac{\sum_{i} [M_{i}]_{pha2}}{\sum_{j} [M_{j}]_{pha1}} = \frac{\left(\frac{\sum_{i} N_{i}}{V}\right)_{pha2}}{\left(\frac{\sum_{j} N_{j}}{V}\right)_{pha1}} = \frac{\left(\frac{A}{\lambda V}\right)_{pha2}}{\left(\frac{A}{\lambda V}\right)_{pha1}} \xrightarrow{\text{same isotope}} D$$

$$= \left(\frac{A_{pha2}}{A_{pha1}}\right) / \underbrace{\left(\frac{V_{pha2}}{V_{pha1}}\right)}_{q} \tag{4}$$

where A is the radioactivity; V, phase volume; N, number of atoms or molecules; α , phase volume ratio.

Sometimes the distribution ratio is expressed in terms of total quantity in moles (n) or mass (m) and termed mass distribution ratio (also called retention factor in chromatography and extraction factor in LLE), D_m , which is correlated to D by eq 5. Since potential confusion may arise between D and D_{m} , the latter variable is employed far less frequently than the

$$D_m = \frac{X_{tot,pha2}}{X_{tot,pha1}} = \alpha D, \ X = m \text{ (mass) or } n \text{ (moles)}$$
(5)

2.2. Separation Performance Evaluation

In many cases, multiple metal elements or compounds are to be separated, so a few parameters are necessary to determine separation efficiency. These include, but are not limited to, recovery, separation factor (SF), enrichment factor (EF), and decontamination factor (DF). Recovery is defined as the ratio of the total quantity (either moles or mass) in phase 2, where the solute is enriched after separation (Q_{pha2}) over the total quantity (Q_0) in the feed before separation $(Q_0 = Q_{pha1} + Q_{pha2})$ the summed quantity in phase 1 and 2, respectively, provided the separating agent used is initially free of the solute to be separated). SF is used to indicate how efficient the separation is between two metal elements (M₁ and M₂). In Co-ppt and LLE, SF can be directly calculated from K or D. However, in IEC, it can be calculated either directly using K or D or indirectly through a reference species for which relative retention (RR) and selectivity over H+ or OH- come into play. RR is the ratio of some quantity (adjusted retention time or volume, D_m , or K) of a species to that of a standard, obtained under identical conditions. Of note, SF is sometimes called selectivity, which is a misnomer. According to IUPAC, selectivity, more precisely selectivity coefficient $(K_{A/B})$, describes the distribution tendency of a target element (M) relative to a reference species (e.g., carrier in Co-ppt $(K_{M/C})$, H^+ in cation exchange chromatography - CEC $(K_{M/(nH)})$, and OH^- in anion exchange chromatography - AEC $(K_{M/(nOH)})$, while SF is the separation between two target elements. EF describes the change in the quantity ratio of two elements from pre- to postseparation and is numerically equal to the reciprocal of DF. A comparison of calculations of these parameters is listed in Table 2.

IEC has a unique way to evaluate the separation performance based on the peak resolution that is defined as R in eq 6, where x is the elution time (t) or the elute volume (V); W_{bb} the peak width at the bottom line (= 4σ = 1.7 FWHM); σ , the peak width at the inflection point; FWHM, the full-width at half-maximum. As a rule of thumb, $R \ge 1.5$ indicates a complete separation and $R \leq 0.8$ denotes no separation. Theoretical plate number and plate height are also parameters that are sometimes useful to evaluate the suitability of the

Table 3. Comparison of Three Isotopes Used in the Experimental Procedures

		S					
	Half-life (yr)	Specific radioactivity (Ci/L)	Matrix	Volume (µL)	Decay mode	Photopeak energy (keV)	Radiation detector
⁶⁰ Co	5.3	0.2	0.01 M HCl or HNO ₃	10	β ⁻ , IT	1173, 1332	Gamma counter ^a
⁶³ Ni	100.1				β^-	N/A	LSC ^b
¹⁵² Eu	13.5				β^- , ϵ	344	Gamma counter ^a

^aEither high purity germanium (HPGe) detector (cooled in liquid N₂) or NaI(Tl) detector. ^bLiquid scintillation counter (LSC).

chromatography column; their use is however out of the scope of this paper:

$$R = \frac{x_{r,A} - x_{r,B}}{\frac{W_{bl,A} + W_{bl,B}}{2}} = \frac{\overbrace{x_{B,\text{max}} - x_{A,\text{max}}}^{\Delta x_{\text{max}}}}{2(\sigma_A + \sigma_B)}$$
(6)

3. EXPERIMENTAL MATERIALS AND APPARATUS

All reagents and devices used are listed in Tables 3 and 4. All chemicals were used as received without further purification. Although purification of chemicals (such as di(2-ethylhexly)-phosphoric acid (HDEHP) used in LLE (vide infra)) is routinely performed in numerous research endeavors, the educational value in this lab course was not lost without this step, as the primary educational objective is to allow students to focus mainly on the concepts, principles, and hands-on experience of these separation techniques. Quantities indicated therein are based on one workstation (most likely a functional hood) for the three experiments and can be scaled up depending on the number of workstations in the classroom. Detailed experiment procedures, exemplary collected data sets, and guidelines of lab reports in the student's handouts can be referred to in the Supporting Information (SI).

4. COPRECIPITATION

Precipitation was one of the first few methods used in radiochemistry by Marie and Pierre Curie to separate Ra in chloride media. However, it is only applicable when the radionuclide to be separated is present in high enough concentrations that exceed its solubility product. For many radioisotopes, the concentration is too low for direct precipitation to pertain. Coprecipitation has been promoted to enrich the target isotopes. Coprecipitation usually requires extraneous chemicals to serve as carriers for the target isotopes (M_1) that await to be separated from others (M_2) (Figure 1). The mechanisms consist of mixed crystallization, surface adsorption, occlusion, and mechanical entrapment. Mixed crystallization usually exhibits the highest selectivity because the nature of isomorphous substitution rigorously requires the carrier and target atoms to be sufficiently similar in size.

In this experiment, separation of 60 Co and 63 Ni was performed via coprecipitation by potassium hexanitrito-cobaltate (also called potassium cobaltinitrite), K_3 Co(NO₂)₆, with stable Co being the isotopic carrier. A batch of Co(NO₃)₂ solutions with different starting concentrations (5, 10, 15, 20, 25, and 30 mM) was prepared and each was spiked with a trace level ($^{\sim}1~\mu$ L, $^{\sim}1-10~n$ Ci) of 60 Co and 63 Ni. The solution was acidified with acetic acid as the masking agent to hold-back 63 Ni¹⁸ and then KNO₂ solution was added to precipitate Co. ¹⁹ The yellow precipitate, K_3 Co(NO₂)₆, forms (eq 7) immediately in concentrated solutions and slowly in diluted solutions. In the classroom demonstration where the concentrations were

dilute, precipitation was accelerated by warming and stirring frequently based on Kahn et al.'s procedure.²⁰ The K₃Co-(NO₂)₆ precipitate was separated from the mother liquor by a vacuum filtration station. Alternatively, phase separation can be done by centrifuging at 3000 rpm for 5 min⁹ in case the precipitate is not well retained on filter paper. ²⁰ The precipitate was washed twice with 5 mL 10 v% acetic acid each time and once with distilled water to remove impurities. Finally, the precipitate was dried using an IR lamp at 110 °C for 5 min, weighed, and counted by the gamma spectrometer for ⁶⁰Co and by the LSC for ⁶³Ni. The filtrate was counted in the same way. The schematic setup is shown in Figure 2 and SI §1.1 provides more details to guide the students. The count rate of each isotope in the solid and in the mother liquor was used to infer the quantity and, further, the distribution ratio, separation factor, and ⁶⁰Co recovery based on equations in Table 2. The results (Table S1, Table S2, and Figure S1 in SI §1.2) showed a trend of the separated ⁶⁰Co as a function of the Co carrier concentration. The method offered the highest recovery of almost 90% with negligible ⁶³Ni in the precipitate. As the starting concentration increased, the recovery decreased because the limiting agent switched from Co(NO₃)₂ to KNO₂, which is to say Co(NO₃)₂ became excess. This is also the reason for the decrease in the distribution ratio of Co. Of note, the factors value remained high since Ni was effectively retained in the liquid phase. It is also possible to examine if the coprecipitation of 60Co follows a homogeneous distribution based on known added quantities and equilibrium quantities of both 60Co and stable Co. In cases where it is difficult or inconvenient to obtain a quantitative recovery, coprecipitation can be applied in conjunction with isotope dilution analysis to obtain more precise quantification since isotope dilution analysis does not require a quantitative recovery of trace analytes:21

$$Co(NO_3)_2 + 7KNO_2 + 2HNO_3$$

 $\to K_3Co(NO_2)_6 \downarrow +NO \uparrow +H_2O + 4KNO_3$ (7)

5. LIQUID-LIQUID EXTRACTION

LLE is one of the most used techniques in separation processes and is preferred in industrial-scale separations, which is ascribed to the ease of changing from batch to multistage operation without significantly increasing the consumption of heat or chemicals. IEC (vide infra), another technique that is prevalent in industrial-scale separations, has this advantage too. In contrast, only a few stages are possible without great expenditure of labor for coprecipitation since crystallization procedures may be by nature slow (long wait-time for precipitate formation) and tedious (with several rinse, dry, and reprecipitation steps).²² Therefore, LLE and IEC outweigh Co-ppt in many applications.

ter. ^{63}Ni is a β^- emitter and does not 2–80% for β^- emitters). All three isotopes negligible impurities that would have

Table 4. Materials and Apparatus Used in the Experimental Procedures

a heterogeneous solution with the cocktail. This has no significant impact on the quantification since tracking the radioactivities of the mother liquor and the filtrate, both of which form homogeneous solutions with the cocktail, yields the accurate radioactivity of the precipitate. Details of principles of used detectors, necessary electronics, and calibrations can be found in Knoll et al.'s work and are ^aHPGe detector used was ORTEC GEM50P4 model equipped with DSPEC-jr 2.0 digital p-ray spectrometer and MAESTRO software. The LSC used was PerkinElmer Packard Tri-Carb model B4430 LSC equipped with QuantaSmart software and the LSC Cocktail was Ultima Gold. The LSC efficiency may be reduced because of the enhanced physical quenching incurred by the precipitate that forms 10 mL malic acid (HOOCCH₂CH(OH)COOH, 0.1 and 0.2 M). not introduced here.

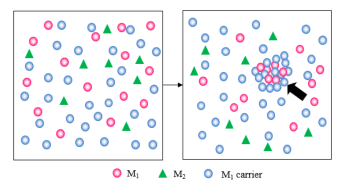


Figure 1. Schematic illustration of the separation of a binary mixture by coprecipitation.

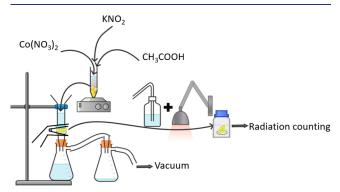


Figure 2. Schematic experimental setup of the separation of ⁶⁰Co from ⁶³Ni by coprecipitation.

An aqueous phase and an ideally immiscible organic phase are mixed. When the liquids settle and phases are separated, solutes may selectively move to one phase more than to the other as driven by the presence of extractants (Figure 3). If there are more than one type of metal ions, either stable or radioactive, they exhibit different distribution behaviors across two phases and thus separation can be achieved.

Extraction experiments were performed under different acidities at 25 $^{\circ}\text{C}$ and consisted of three steps, as listed below and depicted in Figure 3.

- Conditioning. Organic phases (500 μ L 0.1 M HDEHP in dodecane) were contacted with fresh 1 M (H/Na)NO₃ solution for 10 min prior to extraction.
- Extraction. An aqueous mixture of three metal nitrate salts in 1 M (H/Na)NO₃ media (each metal has 0.01 M concentration and 1 μ Ci radioactivity per microtube)

was contacted with an equal volume (500 μ L) of conditioned organic phase in a 1.5 mL screw-cap centrifuge microtube. All microtubes were vigorously shaken for 1 min to achieve thorough mixing before being centrifuged at 3000 rpm for about 3 min to separate the phases. The extraction reaction proceeded as eq 8:

$$Eu(NO_3)_{3(aq)} + 3HDEHP_{(org)}$$

$$\rightleftharpoons Eu(DEHP)_{3(org)} + 3HNO_{3(aq)}$$
(8)

• Counting. A total of 400 µL of the organic phase from the centrifuged microtube was transferred into a labeled flip-top microtube housed in a plastic scintillation vial for radiation counting. The aqueous phase followed the same procedure. A more detailed, step-by-step procedure is provided in SI §2.1 to guide the students. The net counts with error in the characteristic peak for all samples were collected and used to calculate the distribution ratio and the fraction extracted. The counting time can be adjusted based on the radioactivity initially spiked and should be consistent for all samples. The aliquot for radiation detection can also be altered if volume correction is performed afterward.

With this experiment setup, Eu was extracted but Co and Ni were not. 23 D and E values for Eu in pH 0–2 usually fall into the range of 0.5–8 and 30–90%, respectively, based on Figure S2 in SI §2.2 and Cassidy et al.'s work. 23 Both parameter values decrease as the acidity increases because HDEHP is a cation exchanger and the extraction reaction is pH-dependent.

6. ION-EXCHANGE CHROMATOGRAPHY

Many metal ions may be separated using cation exchangers (CEC) since metal ions are inherently cations. However, the distribution behaviors of certain cations of similar chemical properties do not vary greatly. Fortunately, with judicious choice of complexing agent, it is possible to form anionic complexes of these metal ions (e.g., MCl_x^{y-} , $\mathrm{M(NO_3)}_x^{y-}$, etc.) and these complex species can be separated by anion exchangers (AEC) if they exhibit different affinities. For the Dowex strong acid and base resins utilized in the experiment (*vide infra*), the reactions follow eqs 9 and 10 for CEC and AEC, respectively. By varying the concentration of the eluents (usually mineral acids), the fractional ion concentration, existing whether as hydrated cations in CEC or as the complexed anions in AEC, can be varied and large changes in the distribution ratios are thereby produced. Feeds laden with

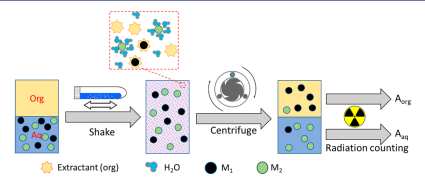


Figure 3. Schematic illustration of the radiochemical separation of a binary mixture by LLE.

a variety of metal ions can thus be eluted at different times or in different volumes and separated (Figure 4). The elution

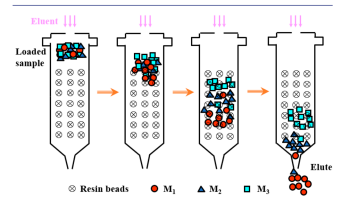


Figure 4. Schematic illustration of the chromatographic separation of a tertiary mixture by IEC with the elution method.

curves of most metallic elements in the periodic table, using hydrochloric acid as the eluent, are shown in the works Modern Nuclear Chemistry²⁴ and Cation-Exchange Techniques in Radiochemistry,²⁵ to which one can refer for in-depth introductions of AEC and CEC, respectively:

CEC:
$$n\overline{Aryl - SO_{3}^{-}H^{+}} + M^{n+}$$

$$\Rightarrow \overline{(Aryl - SO_{3}^{-})_{n}M^{n+}} + nH^{+}$$
(9)
$$AEC: n\overline{Aryl - CH_{2}N(CH_{3})_{3}Cl^{-}} + ML_{x}^{n-}$$

$$\Rightarrow \overline{[Aryl - CH_{2}N(CH_{3})_{3}]_{n}ML_{x}^{n-}} + nCl^{-}$$
(10)

Although CEC and AEC differ in used resins, eluents, mechanisms, and so on (Table 5), separations via IEC generally consist of the following four steps, as listed below and displayed in Figure 5.

 Column preparation. A small wad of glass wool was added to the bottom of a glass column that was then rinsed with appropriate solutions. A slurry of preconditioned ion-exchange resin (strong acid and base used

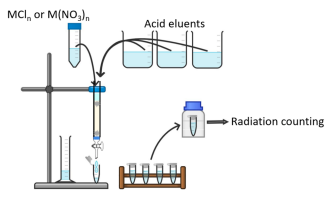


Figure 5. Schematic experimental setup of an anion exchange column used in the chromatographic separation.

here for CEC and AEC, respectively) was placed in the column until reaching about 70–90% of the column height. Another small wad of glass wool was added to the top of the resin bed. The column was rinsed again to even the resin bed and remove the air bubbles or channels. The air pressure was applied carefully (to avoid the risk of contamination) to speed up the dripping and later eluting processes. The drop size from the column was determined by collecting 20 drops in a graduated cylinder. The free column volume (FCV) was determined by rinsing the column with DI water to expel all Cl $^-$ ions, applying a small quantity of NH₄Cl, and eluting the acid with a AgNO $_3$ solution under the column tip. The number of drops when the white AgCl precipitate just appeared was counted as the FCV.

- Loading. The solution mixture containing ⁶³Ni, ⁶⁰Co, and ¹⁵²Eu was loaded onto the resin bed by multiple small additions so as not to put the radioactivity high up on the column reservoir. Fractions of 20 drops were collected in a centrifuge microtube, immediately after loading the feed.
- Elution. CEC and AEC varied greatly in this step due to their different separation mechanisms. CEC followed Dadone et al.'s procedure²⁶ while AEC followed an

Table 5. Comparison of Experimental Conditions for the Separation of ⁵⁹Ni, ⁶⁰Co, and ¹⁵²Eu

			OF C			ATIC		
Experiment setup			CEC			AEC		
Ion exchange resin type		AG 50W-X4 (1 \times 15 cm) or Dowex 50WX8 resin (200–400 mesh) (1.2 \times 22 cm)				AG 1-X4 (1 × 15 cm) or Dowex 1X8 resin (100–200 mesh) (0.29 × 26 cm)		
Resin soaked in		3 M NH ₄ Cl			9 M HCl	9 M HCl		
Column rinsing using		3 M NH ₄ Cl			9 M HCl	9 M HCl		
Loading	Solute	MCl _n or M	MCl_n or $M(NO_3)_n$ (M = Co, Ni, and Eu)					
	Medium	0.1 M malic acid			12 M HCl			
Eluent 1	Acidity	0.2 M malic acid, pH 3.75 for 152Eu			9 M HCl f	9 M HCl for ⁶³ Ni, ¹⁵² Eu		
	Volume ^a	3–8 FCVs			~3 FCVs	~3 FCVs		
Eluent 2 Acidity		0.2 M mali	0.2 M malic acid, pH 4 for ⁵⁹ Ni			4 M HCl for ⁶⁰ Co		
	Volume ^a	3-8 FCVs			~6 FCVs	∼6 FCVs		
Eluent 3	Acidity	0.2 M malic acid, pH 4.25 for 60Co			N/A			
	Volume ^a	3–8 FCVs						
Results		⁶⁰ Co	⁶³ Ni	¹⁵² Eu	⁶⁰ Co	⁶³ Ni	¹⁵² Eu	
Eluent 1	$D \left(mL/g \right)^{b}$	~32	~8	unretained	~40	unretained		
Eluent 2		~14	~14 unretained		unretained			
Eluent 3		unretained	unretained			N/A		

^aExact eluting volume is subject to the column size. ^bD values by CEC are from Dadone et al.'s work. ²⁶ and those by AEC are from Kraus et al.'s work. ²⁷

established procedure used in our classroom settings (SI §3.1.1). For both methods, eluents with different acidities (Table 5) were added to elute different isotopes and meanwhile the fraction collection was continued. The air pressure was applied to the column and the volumetric flow rate of the mobile phase was ~ 3 mL/min.

Counting. The count rates of ⁵⁹Ni, ⁶⁰Co, and ¹⁵²Eu in the collected elute fractions, in the sample solution before and after loading, and in the air-dried column upon the completion of fraction collection, were accomplished in the same way as mentioned above.

The D values are summarized in Table 5, from which we can see the separation among the three radioisotopes was effortlessly achieved by CEC. This terrific separation performance was also evidenced by the well-resolved elution bands of the three isotopes with a peak resolution of ~2.3 between 60 Co and 63 Ni. In contrast, AEC did not differ between 63 Ni and 152 Eu. Other parameters in assessing separation performance, such as the elution peak resolution, can then be estimated as well.

The actual practice in CHEM/NE C146 used a different isotope trio to teach students about other isotopes. Comprehensive details on each eluted fraction and the elution profile are in Table S4 and Figure S3 in SI §3.1.2, from which the aforementioned thermodynamic parameters can be calculated. The exact procedure and analysis can be made on the proposed ⁵⁹Ni, ⁶⁰Co, and ¹⁵²Eu trio, which is used in the other two proposed separation experiments. Using the same isotopes throughout all experiments offers easier safety control, fewer hazards, and lower cost than introducing new ones, however, at the expense of limited knowledge transfer to students. Educators may tailor to their own needs and make adjustments in selecting appropriate isotopes. For circumstances where experiment time and space layout are constrained, educators may consider using miniaturized isotope generators ^{28,29} and bypassing the column preparation and metal solution loading steps. A ¹³⁷Cs/^{137m}Ba isotope generator was used in our practice and elsewhere ³⁰ to demonstrate CEC and radioactive decay (SI §3.2) since the students had already learned about preseparation operations in the previous AEC experiment. This device is typically used to determine the halflife and decay constant, which are beyond the scope of this article. Those who are interested in this subject may consult SI §3.2 and a variety of sources elsewhere.

7. CONCLUSIONS

The principles of three separation techniques were introduced along with a few fundamental thermodynamic concepts frequently used in the separation process. Applications of these techniques to the separations of a rare earth element (Eu) and two transition metals (Co and Ni) were proposed in a way that is feasible for N&RC-relevant laboratory sessions and/or courses. The introduced techniques are not limited to radionuclide separations. They have a wide range of applications where metal recovery is sought, such as the reprocessing of electronic waste. For institutions that do not easily have access to radioisotopes, these techniques can still be instructed since the quantification by radiation detectors can be substituted by ICP-MS or ICP-AES at the expense of complicated sample dilution and more liquid waste generation. Our teaching experience over the past few years has

demonstrated that students with a background in nuclear engineering generally perform radiochemical separation experiments less well than those with a background in chemistry and/or chemical engineering. In contrast, chemistry and/or chemical students typically have more difficulties in radiation detection of radioactive samples than nuclear engineering students (Figure 6). The radiochemical separation experiments

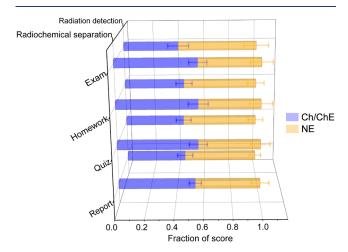


Figure 6. Comparison of the course performance of 20 students majoring in chemistry/chemical engineering (Ch/ChE) or nuclear engineering (NE). The score is normalized to 1, and a value of 0.5 indicates an equal score between Ch/ChE and NE students.

discussed here can serve as a bridge between the two domains. Of note, the techniques discussed in this work by no means cover all areas of N&RC but are substantial parts of radiochemical separation. By taking advantage of radiation detection courses, N&RC courses and pertinent programs can advance and meet the deficiencies in both education and industrial settings.

HAZARDS

Potential Health Hazards Encountered Are As Follows

Radiological Hazards. All isotopes used in this study, ⁶⁰Co, ⁶³Ni, and ¹⁵²Eu, are radioactive materials and should be handled only in specifically designated facilities in accordance with appropriate safety controls. The National Institute for Occupational Safety and Health (NIOSH) Hierarchy of Controls³¹ was closely adhered to in order to prevent the spreading of radioactivity throughout the working area, the release of airborne radioactivity outside of the hood, and the potential contamination of the students. For the duration of all experiments, the drains were covered with landscaping fabric or similar materials, and the workbenches were covered with diaper paper. All chemicals and tools were placed in polypropylene trays lined with diaper paper. The quantity handled by the students was restricted to the bare minimum necessary to ensure that the statistical character of radioactive decay had the least possible impact on counting errors. In the LLE experiment, screw-cap vials were used to substitute the conventional large and leak-prone separatory funnels. In the precipitation experiment, a planchet held with forceps was practiced to keep a safe distance between the students and the radioactive substance. We suggest that students work in pairs using a "hot hands/cold hands" approach. One student ("hot hands") works directly with radioactive materials and is

double-gloved at all times. The second student ("cold hands") performs nonradioactive work to assist his or her partner, such that no contamination is spread to clean regions. Students and graduate student instructors should conduct full swipes and surveys of the workstations and surrounding floor after each lab experiment to either ensure no contamination or immediately identify contamination.

Chemical Hazards. Mineral acids (HNO₃, HCl), organic acids (acetic acid, malic acid, and HDEHP), and organic solvents (kerosene) must be handled in a properly ventilated hood in accordance with standard operating procedures.

Physical Hazards. Radiation detection equipment poses ergonomic and electrical hazards and should be used by proper guidance.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available at https://pubs.acs.org/doi/10.1021/acs.jchemed.2c01193.

Experimental details, lab handouts, and exemplary gathered data (PDF, DOCX)

AUTHOR INFORMATION

Corresponding Author

Yufei Wang — Department of Nuclear Engineering, University of California, Berkeley, California 94709, United States; Chemical Sciences Division, Lawrence Berkeley National Laboratory, Berkeley, California 94720, United States; orcid.org/0000-0002-5780-112X; Email: yufeiwang@berkeley.edu

Author

Rebecca J. Abergel — Department of Nuclear Engineering, University of California, Berkeley, California 94709, United States; Chemical Sciences Division, Lawrence Berkeley National Laboratory, Berkeley, California 94720, United States; Oorcid.org/0000-0002-3906-8761

Complete contact information is available at: https://pubs.acs.org/10.1021/acs.jchemed.2c01193

Author Contributions

Yufei Wang: Conceptualization, Methodology, Investigation, Formal analysis, Visualization, Software, Writing—original draft, review and editing. Rebecca J. Abergel: Supervision, Resources, Writing—review and editing, Funding acquisition.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

We are grateful to graduate student instructors who have taught the Nuclear Chemistry and Radiochemistry laboratory class at UC Berkeley over the past few years, Drs. J. Shusterman, E. May, E. Uribe, and K. Shield. Also, thanks to Hailie M.T. O'Bryan for filming the experiments for remote learning during the COVID-19 pandemic. The assembly of this manuscript was supported by the U.S. Department of Energy, Office of Nuclear Energy's Nuclear Energy University Programs (DE-NE0008949).

ABBREVIATIONS

Co-ppt coprecipitation.

DF decontamination factor.

EF enrichment factor.

FCV free column volume.

FWHM full width at half-maximum.

HDEHP also called D(2)EHPA,di(2-ethylhexly)phosphoric acid.

HPGe high purity germanium.

I/A/CEC ion/anion/cation exchange chromatography.

ICP-MS/AES inductively coupled plasma mass spectrome-

try/atomic emission spectroscopy LLE liquid—liquid extraction.

LSC liquid scintillation counter.

N&RC nuclear chemistry and radiochemistry.

NIOSH National Institute for Occupational Safety and

RR relative retention.

SF separation factor.

UC Berkeley University of California at Berkeley.

REFERENCES

- (1) Pernaa, J.; Law, G. T. W.; Ranjan, S. The Relevance of Radiochemistry: Perceptions of Future Radiochemists. *J. Chem. Educ.* **2021**, *98*, 8.
- (2) Assuring a Future U.S.-Based Nuclear and Radiochemistry Expertise; National Academies Press: Washington, D.C., 2012; p 13308. DOI: 10.17226/13308.
- (3) Shi, W.-Q.; Zhao, Y.-L.; Chai, Z.-F. Nuclear and Radiochemistry in China: Present Status and Future Perspectives. *Radiochim. Acta* **2012**, *100* (8–9), 529–539.
- (4) Wang, Y.-H.; Zhang, G.-H.; Xiang, Y.-Q.; Yuan, W.-L.; Fu, J.; Wang, S.-L.; Xiong, Z.-X.; Zhang, M.-D.; He, L.; Tao, G.-H. Virtual Reality Assisted General Education of Nuclear Chemistry and Radiochemistry. *J. Chem. Educ.* **2022**, *99* (2), 777–786.
- (5) Crosier, J. K.; Cobb, S. V. G.; Wilson, J. R. Experimental Comparison of Virtual Reality with Traditional Teaching Methods for Teaching Radioactivity. *Education and Information Technologies* **2000**, 5 (4), 329–343.
- (6) White, A. E. Cloud Chamber Kit for Active Learning in a First-Year Undergraduate Nuclear Science Seminar Class. *Physics Education* **2022**, *57* (4), 045034.
- (7) Den Auwer, C.; Bayle, S.; Beccia, M. R.; Bosio, S.; Creff, G.; Jeanson, A.; Michel, H.; Pitiot, C.; Zurita, C. A Table-Top Cloud Chamber to Observe Radioactivity in the Uranium Decay Chain. *J. Chem. Educ.* **2022**, *99* (2), 646–653.
- (8) Holgersson, S. Hands-On Model of the Principle of Isotope Dilution Analysis for Use in an Interactive Teaching and Learning Classroom Exercise. *J. Chem. Educ.* **2021**, *98* (4), 1208–1220.
- (9) Young, J. D.; George, E. P. An Isotopic Method for Estimation of Urinary Potassium. *J. Nucl. Med.* **1965**, *6* (4), 265.
- (10) Livermore, A. H.; Scott, A. F. Radiochemistry—Advanced Undergraduate Work. *J. Chem. Educ.* **1960**, 37 (8), 387.
- (11) Coryell, C. D.; Irvine, J. W., Jr. Teaching Radiochemistry at Massachusetts Institute of Technology. *J. Chem. Educ.* **1951**, 28 (1), 14.
- (12) Pacer, R. A.; Ehmann, W. D.; Yates, S. W. An Isotope Dilution Analysis Experiment with Phase Isolation by Electrodeposition. *J. Chem. Educ.* **1989**, *66* (7), 603.
- (13) Williams, K. R.; Lipford, L. C. Isotropic Dilution Analysis and Secular Equilibrium Study: Two Complementary Radiochemistry Experiments. *J. Chem. Educ.* **1985**, *62* (10), 894.
- (14) Voige, W. H. Extraction and Assay of Ornithine Decarboxylase: A Laboratory Experiment That Introduces Principles of Radiochemical Enzymatic Assay. *J. Chem. Educ.* **1997**, *74* (8), 988.

- (15) Wang, C. H.; Willis, D. L.; Loveland, W. D. Radiotracer Methodology in the Biological, Environmental, and Physical Sciences; Prentice-Hall, 1975.
- (16) Knoll, G. F. Radiation Detection and Measurement; John Wiley & Sons, 2010.
- (17) Marie Curie Research Breakthroughs (1897–1904); American Institute of Physics, 2011. https://web.archive.org/web/20111118025917/http://www.aip.org/history/curie/resbr2.htm (accessed 06-20-2021).
- (18) Vogel, A. I. Vogel's Textbook of Macro and Semimicro Qualitative Inorganic Analysis, 5th ed.; Longman: London, 1979.
- (19) Bate, L. C.; Leddicotte, G. W. THE RADIOCHEMISTRY OF COBALT 1961, NAS-NS-3041-4842901.
- (20) Kahn, B. Procedures for the Analysis of Some Radionuclides Absorbed on Soil; Oak Ridge National Lab., 1955.
- (21) Inagaki, K.; Zhu, Y.; Miura, T.; Chiba, K. Coprecipitation in Trace Element Analysis. In *Encyclopedia of Analytical Chemistry*; Meyers, R. A., Ed.; John Wiley & Sons, Ltd: Chichester, UK, 2009; p a9065. DOI: 10.1002/9780470027318.a9065.
- (22) Tipson, R. S. Theory, Scope, and Methods of Recrystallization. *Anal. Chem.* **1950**, 22 (5), 628–636.
- (23) Cassidy, P. W. Solvent Extraction of Europium, Cobalt and Nickel with DI-2-Ethylhexyl Phosphate. 1970.
- (24) Loveland, W. D.; Morrissey, D. J.; Seaborg, G. T. Modern Nuclear Chemistry; Wiley, 2017.
- (25) Massart, D. L. CATION-EXCHANGE TECHNIQUES IN RADIOCHEMISTRY. 1971, DOI: 10.2172/4699019.
- (26) Dadone, A.; Baffi, F.; Frache, R.; Cosma, B.; Mazzucotelli, A. Organic Acid Solutions in the Chromatography of Inorganic Ions VIII. Cation-Exchange of Mn(II), Cd(II), Co(II), Ni(II), Zn(II), Cu(II), Fe(III), Sc(III), Y(III), Eu(III), Dy(III), Ho(III), Yb(III), Ti(IV) and Nb(V) in Malate Media. *Chromatographia* 1981, 14 (1), 32–38.
- (27) Kraus, K. A.; Nelson, F. Anion Exchange Studies of the Fission Products. Proceedings International Conference on the Peaceful Uses of Atomic Energy 1955, VII, 837.
- (28) Pinajian, J. J. A Cesium-137-Barium-137m Isotope Generator. *J. Chem. Educ.* **1967**, *44* (4), 212.
- (29) Crater, H. L.; Macchione, J. B.; Gemmill, W. J.; Kramer, H. H. Radioisotope generators for introductory laboratory use. *J. Chem. Educ.* **1969**, *46*, 287 DOI: 10.1021/ed046p287.
- (30) Konkankit, C. C.; Marker, S. C.; Bigham, N. P.; Dale, D. S.; Zax, D. B.; Lorey, D. R., II; Wilson, J. J. Development and Implementation of Nuclear Chemistry Experiments at the Undergraduate Level. *J. Chem. Educ.* **2021**, *98* (12), 3831–3840.
- (31) Centers for Disease Control and Prevention (USA). National Institute for Occupational Safety and Health (NIOSH): Hierarchy of controls, 2023. https://www.cdc.gov/niosh/topics/hierarchy.

□ Recommended by ACS

Highly Efficient Extraction Chromatography Resin Containing Hexa-n-Octyl Nitrilotriacetamide (HONTA) for Selective Recovery of Plutonium from Acidic Feeds

Rajesh B. Gujar, Willem Verboom, et al.

MARCH 31, 2023

INDUSTRIAL & ENGINEERING CHEMISTRY RESEARCH

READ 🗹

CeRES Process—Separation of Cerium from Lanthanum by Redox Extraction and Stripping

Yujian Zhou, Jan J. Weigand, et al.

NOVEMBER 22, 2022

ACS SUSTAINABLE CHEMISTRY & ENGINEERING

READ 🗹

Simple, Fast, and Selective Dissolution of $\mathrm{Eu_2O_3}$ in an Ionic Liquid as a Sustainable Paradigm for Lanthanide–Actinide Separations in Radioactive Waste Remediation

Parveen Kumar Verma, Sk. Musharaf Ali, et al.

DECEMBER 18, 2022

INORGANIC CHEMISTRY

READ 🗹

Size Selective Ligand Tug of War Strategy to Separate Rare Earth Elements

Katherine R. Johnson, Santa Jansone-Popova, et al

JANUARY 25, 2023

JACS AU

READ 🗹

Get More Suggestions >